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(54) Title: INTERFERON-ALPHA 54

Net Ala Leu Pro Phe Val Leu Lau Het Ala Leu Val Val Leu Asn Cys Lye Ser Ile Cys Arg GCC TTG CCC TTT GTT TTA CTG ATG GCC CTG GTG GTG CTC AAC TGC AAG TCA ATC TGT Ser Led Gly Cys Asp Led Pro Gln The His Ser Led Ser Asn Arg Arg The Led Net Ile TCT CTG GGC TGT GAT CTG CCT CAG ACC CAC AGC CTG AGT AAC AGG AGG ACT TTG ATG ATA Net Ale Gin Het Gly Ard Ile Ser Pro Phe Ser Cys Leu Lys Asp Ard Sis Asp Phe Gly Ard GCA CAA ARG GCA CAT GAC TIT GGA The tro cln clu clu the Asp cly Asp cln the Cin Lys Ala cin Ala Ile Set Val Lee TTT CCT CAG CAG CAG TTT CAT CCC AAC CAG TTC CAG AAG CCT CAA CCC ATC TCT GTC CTC Bis Glu Het lie Gln Gln Thr Phe Asn Leu Phe Ser Thr Lys Asp Ser Ser Ale Thr Trp CAT GAG AND ATC CAG CAG ACC TTC AAT CTC TTC AGC ACA AAG GAC TCA TCT GCT ACT TGG Asp Glo The Lea Lea Asp Lys Phe Tyr The Glo Lea Tyr Glo Glo Lea Aso Asp Lea Glo GAT GAG ACA CIT CIA GAC AAA TIC TAC ACT GAA CIT TAC CAG CAG CIG AAT GAC CIG GAA Ala Cys Het Het Gin Glu Val Gly Val Glu Asp Thr Pro Leu Het Asn Val Asp Ser Ile GCC TGT ATG ATG CAG GAG GTT GGA GTG GAA GAC ACT CCT CTG ATG AAT GTG GAC TCT ATC Let The Val Arg Lys Tyr Phe Gln Arg Ile The Let Tyr Let The Glt Lys Lys Tyr Ser CTG ACT GTG AGA AAA TAC TTT CAA AGA ATC ACT CTC TAT CTG ACA GAG AAG AAA TAC AGC Fro Cys Ale Trp Glu Vel Vel Arg Ale Glu Ile Het Arg Ber Phe Ser Leu Ser Ale Asn CCT TGT GCA TGG GAG GTT GTC AGA GCA GAA ATC ATG AGA TCC TTC TCT TTA TCA GCA AAC Leu Gin Giu Arg Leu Arg Arg Lye Giu TTG CAA GAA AGA TTA AGG AGG AAG GAA

(57) Abstract

New polypeptide, called IFN-α54, produced by E. coli transformed with a newly isolated and characterized human IFN-a gene. The polypeptide exhibits interferon activities such as antiviral activity, cell growth regulation, and regulation of production of cell-produced substances.

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## INTERFERON-ALPHA 54

## Description

## Technical Field

The invention is in the field of biotech-5 nology. More particularly it relates to a polypeptide having interferon (IFN) activity, DNA that codes for the polypeptide, a recombinant vector that includes the DNA, a host organism transformed with the recombinant vector that produces the polypeptide, pharma-. 10 ceutical compositions containing the polypeptide, and therapeutic methods employing the polypeptide.

Background Art IFNs are proteins with antiviral, immunomodulatory, and antiproliferative activities produced 15 by mammalian cells in response to a variety of inducers (see Stewart, W.E., The Interferon System, Springer-Verlag, New York, 1979). The activity of IFN is largely species specific (Colby, C., and Morgan, M. J., Ann. Rev. Microbiol. 25:333-360 (1971) and thus 20 only human IFN can be used for human clinical studies. Human IFNs are classified into three groups,  $\alpha$ ,  $\beta$ , and  $\gamma$ , (Nature, 286:110, (1980)). The human IFN- $\alpha$  genes compose a multigene family sharing 85%-95% sequence homology (Goeddel, D. V., et al, Nature 290:20-27 25 (1981) Nagata, S., et al, J. Interferon Research 1:333-336 (1981)). Several of the IFN- $\alpha$  genes have

been cloned and expressed in E.coli (Nagata, S., et



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al, Nature 284:316-320 (1980); Goeddel, D. V., et al,
Nature 287:411-415 (1980); Yelverton, E., et al,
Nucleic Acids Research, 9:731-741, (1981); Streuli,
M., et al, Proc Nat Acad Sci (USA), 78:2848-2852. The
resulting polypeptides have been purified and tested
for biological activities associated with partially
purified native human IFNs and found to possess similar activities. Accordingly such polypeptides are
potentially useful as antiviral, immunomodulatory, or
antiproliferative agents.

A principal object of the present invention is to provide a polypeptide having interferon activity that is produced by an organism transformed with a newly isolated and newly characterized IFN-α gene.

15 This polypeptide is sometimes referred to herein as IFN-α54. Other objects of the invention are directed to providing the compositions and hosts that are used to produce this polypeptide and to therapeutic compositions and methods that use this polypeptide as an active ingredient.

## Disclosure of the Invention

One aspect of the invention is a polypeptide having interferon activity and comprising the amino acid sequence:

CysAspLeuProGln ThrHisSerLeuGly HisArgArgThrMet MetLeuLeuAlaGln MetArgArgIleSer LeuPheSerCysLeu LysAspArgHisAsp PheArgPheProGln GluGluPheAspGly AsnGlnPheGlnLys AlaGluAlaIleSer ValLeuHisGluVal IleGlnGlnThrPhe AsnLeuPheSerThr LysAspSerSerVal AlaTrpAspGluArg LeuLeuAspLysLeu TyrThrGluLeuTyr GlnGlnLeuAsnAsp LeuGluAlaCysVal MetGlnGluValTrp ValGlyGlyThrPro LeuMetAsnGluAsp SerIleLeuAlaVal ArgLysTyrPheGln ArgIleThrLeuTyr LeuThrGluLysLys TyrSerProCysAla TrpGluValValArg AlaGluIleMetArg SerPheSerSerSer ArgAsnLeuGlnGlu ArgLeuArgArgLys Glu



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A second aspect of the invention is a DNA unit or fragment comprising a nucleotide sequence that encodes the above described polypeptide.

A third aspect of the invention is a cloning 5 vehicle or vector that includes the above described DNA.

A fourth aspect of the invention is a host that is transformed with the above described cloning vehicle and that produces the above described polypeptide.

A fifth aspect of the invention is a process for producing the above described polypeptide comprising cultivating said transformed host and collecting the polypeptide from the resulting culture.

Another aspect of the invention is a pharmaceutical composition having interferon activity comprising an effective amount of the above described polypeptide admixed with a pharmaceutically acceptable carrier.

Still another aspect of the invention is a method of providing interferon therapy to a human comprising administering a therapeutically effective amount of the above described polypeptide to the human.

# 25 Brief Description of the Drawings

Figure 1 is a partial restriction map which shows the two XhoII restriction sites that produce a homologous 260 base pair DNA fragment from the IFN- $\alpha$ l and IFN- $\alpha$ 2 structural genes. Data for this map are from Streuli, M., et al Science, 209:1343-1347 (1980).

Figure 2 depicts the sequencing strategy used to obtain the complete DNA sequence of the IFN- $\alpha$ 54 gene coding region. Bacteriophage mp7: $\alpha$ 54-1



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DNA served as the template for sequences obtained with primers A, H and F and bacteriophage mp7: a54-2 DNA was the template for sequences obtained with primers E and G. The crosshatched area of the gene depicts the region that encodes the 23 amino acid signal peptide and the open box depicts the region that encodes the mature polypeptide. The scale, in base pairs, is numbered with 0 representing the ATG start codon of preinterferon. The arrows indicate the direction and extent of sequencing with each primer.

Figure 3 is the nucleotide sequence of the structural gene coding for IFN-a54 including some of the flanking 5'- and 3'- noncoding regions of the gene. The region coding for preinterferon and the 15 mature polypeptide begins with the ATG codon at position 61 and terminates with the TAA codon at position 628.

Figure 4 is a partial restriction map of the coding region of the IFN-\$\alpha\$54 gene. The crosshatching represents the region that encodes the 23 amino acid signal peptide and the open box represents the gene coding sequence for the mature polypeptide. The scale, in base pairs, is numbered with 0 representing the ATG start codon of preinterferon.

Figure 5 shows the amino acid sequence of the 23 amino acid signal polypeptide and the 166 amino acid mature IFN-α54 coded for by the gene depicted in Figure 3. The 189 amino acid sequence is displayed above the corresponding nucleotide sequence. Amino acid 24, cysteine, is the first amino acid of the mature IFN-α54 protein.

Figure 6 is the DNA sequence of the <u>E. coli</u> trp promoter and the gene of Figure 3 which was inserted between the <u>EcoRI</u> and <u>PvuII</u> sites of the



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plasmid pBWll. The amino acid sequence of Figure 5 is written above the corresponding DNA sequence and the location of the restriction sites used in the construction of the expression plasmid are indicated.

Figure 7 is a diagram of the expression plasmid, pCS12.

# Modes for Carrying Out the Invention

In general terms IFN- $\alpha$ 54 was made by identifying and isolating the IFN- $\alpha$ 54 gene by screening a library of human genomic DNA with an appropriate IFN- $\alpha$ 50 DNA probe, constructing a vector containing the IFN- $\alpha$ 54 gene, transforming microorganisms with the vector, cultivating transformants that express IFN- $\alpha$ 54 and collecting IFN- $\alpha$ 54 from the culture. A preferred embodiment of this procedure is described below.

## DNA Probe Preparation

17:517-526 (1979).

Total cytoplasmic RNA was extracted from human lymphoblastoid cells, Namalwa, which had been induced for IFN production by pretreatment with 20 5-bromodeoxyuridine and Newcastle Disease Virus (NDV). The poly(A) (polyadenylic acid)-containing messenger RNA (mRNA) was isolated from total RNA by chromatography on oligo(dT)-cellulose (type 3 from Collaborative Research; Aviv, H., and Leder, P., Proc 25 Natl Acad Sci (USA), 69:1408-1412, (1972)) and enriched for IFN mRNA by density gradient centrifugation on 5%-20% sucrose gradients. Fractions containing IFN mRNA were identified by translating the mRNA by microinjecting aliquots of each fraction into 30 Xenopus oocytes and determining the IFN activity of the products of the translations according to a method described by Colman, A., and Morser, J., Cell,

The Namalwa cell human IFN enriched mRNA was used to construct complementary DNA (cDNA) clones in E. coli by the G/C tailing method using the PstI site of the cloning vector pBR322 (Bolivar, F., et al,

5 Gene, 2:95-113 (1977)). A population of transformants containing approximately 50,000 individual cDNA clones was grown in one liter of medium overnight and the total plasmid DNA was isolated therefrom.

The sequences of two IFN-α clones (IFN-αl and IFN-α2) have been published (Streuli, M., et al, Science, 209:1343-1347 (1980)). Examination of the DNA sequences of these two clones revealed that the restriction enzyme XhoII would excise a 260 bp fragment from either the IFN-αl or the IFN-α2 gene (see Figure 1). XhoII was prepared in accordance with the process described by Gingeras, T.R., and Roberts, R.J., J Mol Biol, 118:113-122 (1978).

One mg of the purified total plasmid DNA preparation was digested with XhoII and the DNA frag-20 ments were separated on a preparative 6% polyacrylamide gel. DNA from the region of the gel corresponding to 260 bp was recovered by electroelution and recloned by ligation into the BamHI site of the single strand bacteriophage ml3:mp7. Thirty-six clones were 25 picked at random and the single stranded DNA isolated therefrom and sequenced. The DNA sequences of four of these clones were homologous to known IFN- $\alpha$  DNA sequences. Clone mp7:  $\alpha$ -260, with a DNA sequence identical to IFN-al DNA (Streuli, M. et al, Science, 30 209:1343-1347 (1980)) was chosen as a highly specific hybridization probe for identifying additional IFN- $\alpha$ DNA sequences. This clone is hereinafter referred to as the "260 probe."



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## Screening of Genomic DNA Library

In order to isolate other IFN- $\alpha$  gene sequences, a 32p-labelled 260 probe was used to screen a library of human genomic DNA by in situ hybridiza-5 tion. The human gene bank, prepared by Lawn, R.M., et al, Cell, 15:1157-1174 (1978), was generated by partial cleavage of fetal human DNA with HaeIII and AluI and cloned into bacteriophage  $\lambda$  Charon 4A with synthetic EcoRI linkers. Approximately 800,000 clones 10 were screened, of which about 160 hybridized with the 260 probe. Each of the 160 clones was further characterized by restriction enzyme mapping and comparison with the published restriction maps of 10 chromosomal IFN genes (Nagata, S., et al, J Interferon Research, 15 1:333-336 (1981)). One of the clones, hybrid phage  $\lambda 4A: \alpha 54$  containing a 16 kb insert, was characterized as follows. A DNA preparation of  $\lambda 4A: \alpha 54$  was cleaved with HindIII, BglII, and EcoRI respectively, the fragments separated on an agarose gel, transferred to a 20 nitrocellulose filter (Southern, E.M., J Mol Biol, 98:503-517 (1977)) and hybridized with 32p-labelled 260 probe. This procedure localized the IFN-a54 gene to a 3.9 kb EcoRI restriction fragment which was then isolated and recloned, in both orientations, by 25 ligation of the fragment into EcoRI cleaved ml3:mp7. The two subclones are designated mp7: $\alpha54-1$  and mp7: $\alpha$ 54-2. The -1 designation indicates that the single-stranded bacteriophage contains insert DNA complementary to the mRNA (the minus strand) and the 30 -2 designation indicates that the insert DNA is the

same sequence as the mRNA (the plus strand).



## Sequencing of the IFN-a54 Gene

The Sanger dideoxy-technique was used to determine the DNA sequence of the IFN-a54 gene. strategy employed is diagrammed in Figure 2, the DNA 5 sequence thus obtained is given in Figure 3, and a partial restriction enzyme map of the IFN- $\alpha54$  gene is illustrated in Figure 4. Unlike many genes from eukaryotic organisms, but analogous to other IFN chromosomal genes which have been characterised 10 (Nagata, S., et al, Nature, 287:401-408 (1980); Lawn, R.M., et al, Science, 212:1159-1162 (1981); Lawn, R.M., et al, <u>Nucleic Acids Res.</u>, 9:1045-1052 (1981); Nagata, S., et al, J Interferon Research, 1:333-336 (1981); Lawn, R.M., et al, Proc Natl Acad Sci (USA), 15 78:5435-5439 (1981)), the DNA sequence of this gene demonstrates that it lacks introns. Homology to protein sequence information from these published IFN- $\alpha$  genes made it possible to determine the correct translational reading frame and thus allowed the 20 entire 166 amino acid sequence of IFN-α54 to be predicted from the DNA sequence as well as a precursor segment, or signal peptide, of 23 amino acids (Figure 5). The DNA sequence of the IFN- $\alpha54$  gene and the amino acid sequence predicted therefrom differ sub-25 stantially from the other known IFN- $\alpha$  DNA and IFN- $\alpha$ amino acid sequences.

# Plasmid Preparation and Host Transformation

Assembly of the plasmid for direct expression of the IFN-a54 gene involved replacing the DNA fragment encoding the 23 amino acid signal polypeptide of preinterferon with a 120 bp <a href="EcoRI/Sau3A">EcoRI/Sau3A</a> promoter fragment (E.coli trp promoter, operator, and trp leader ribosome binding site preceding an ATG initia-



tion codon) and using the naturally occurring HincII site, 139 bp 3'- of the TAA translational stop codon, to insert the gene into a cloning vehicle derived from the plasmid pBR322. The complete DNA sequence of the promoter and gene fragments inserted between the EcoRI and PvuII sites of pBR322 is shown in Figure 6 which also shows the exact location of relevant cloning sites. Details of the construction are described below.

The coding region for mature IFN-\$\alpha\$54 encompasses a Sau3A site between codons for amino acids 2 and 3 and an XbaI site between codons for amino acids 81 and 83. The 239 bp Sau3A to XbaI fragment was isolated on a 6% polyacrylamide gel following a

Sau3A/XbaI double-digest of the 3.9 kb EcoRI genomic fragment. This fragment was ligated to the 120 bp EcoRI/Sau 3A promoter fragment. The promoter fragment contained a synthetic HindIII restriction site, ATG initiation codon, the initial cysteine codon (TGT) and

a <u>Sau</u> 3A "sticky end". The ligation mixture was digested with <u>EcoRI</u> and <u>XbaI</u> to enrich for the desired product and ligated with an <u>EcoRI/XbaI</u> digested vector fragment pBW11 (derived from pBR322 which contained unique <u>EcoRI</u> and <u>XbaI</u> restriction sites). The

ligation mixture was used to transform E.coli MM294 (Backman, K., et al, Proc Natl Acad Sci (USA), 73:4174-4178 (1976)). The desired correct transformation product, designated pCS10, was identified by restriction enzyme mapping. DNA from this interme-

diate plasmid was prepared, digested with XbaI and PvuII, and the large fragment containing the promoter and the 5'-portion of the gene was used as a vector for reconstituting the 3'-end of the gene. Referring to the restriction enzyme sites shown in Figure 6, the



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397 base pair Xba to HincII fragment encompassing the 3'-codon region, the TAA translational stop codon and an additional 139 base pairs of 3'-noncoding sequence was isolated on a 4% polyacrylamide gel following an 5 XbaI/HincII double-digest of the 3.9 kb genomic EcoRI fragment. This XbaI to HincII fragment was ligated to the Xba/PvuII digested pCS10, the ligation mixture was used to transform E.coli MM294, and correct transformants (3 out of 100 screened) were identified by restriction enzyme mapping. Figure 7 is a diagram of one of the final expression constructs obtained, which is designated pCS12. Other prokaryotic hosts such as bacteria other than E.coli may, of course, be transformed with this or other suitable constructs to replicate the IFN-α54 gene and/or to produce IFN-α54.

### Cultivation of Transformants

Bacteria transformed with the IFN-α54 gene may be cultivated in an appropriate growth medium, such as a minimum essential medium, that satisfies the nutritional and other requirements needed to permit the bacteria to grow and produce IFN-α54. If the bacteria are such that the protein is contained in their cytoplasm, the IFN-α54 may be extracted from the cells by lysing the cells such as by sonication and/or treatment with a strong anionic solubilizing agent such as sodium dodecyl sulfate. Further purification of the extract may be achieved by affinity chromatography, electrophoresis, or other protein purification techniques.

IFN- $\alpha$ 54 produced in accordance with the invention is believed to be distinct from the corresponding native protein in several respects.



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Firstly, because the IFN- $\alpha$ 54 gene was expressed by bacterial hosts that utilize N-formyl-methionine and/or methionine to initiate translation, some or all of the bacterially produced IFN-a54 molecules are pre-5 ceded by an N-formyl-methionine or methionine group. Some of the N-formyl-methionine or methionine groups could be removed by natural in vivo bacterial cleavage mechanisms. This would result in a mixture of molecules, some of which would include an initial N-10 formyl-methionine or methionine and others that would not. All such IFN- $\alpha$ 54 molecules, those containing an initial N-formyl-methionine or methionine, those not containing an N-formyl-methionine or methionine and any mixture thereof, are encompassed by the present 15 invention. Secondly, the amino acid residues of the bacterially produced polypeptide are unsubstituted whereas the residues of the native protein may be substituted with sugar groups, ACTH or other moie-Also, native IFN- $\alpha$  extracts consist of mixtures 20 of various IFN molecules whereas the bacterially produced IFN- $\alpha$ 54 is homogeneous; that is, bacterially produced IFN-a54 does not contain functionally related polypeptides. Accordingly, the invention contemplates producing IFN-α54-containing compositions having bio-25 logical activity that is attributable solely to IFN- $\alpha$ 54 and/or said terminal N-formyl-methionine or methionine derivatives thereof.

## Biological Testing of IFN- $\alpha$ 54

IFN-α54-containing cell sonicates were

30 tested in vitro and found to have the following
activities: (1) inhibition of viral replication of
vesicular stomatitis virus (VSV) and herpes simplex
virus-1 (HSV-1); (2) inhibition of tumor cell growth;



(3) inhibition of colony formation by tumor cells in soft agar; (4) activation of natural killer (NK) cells; (5) enhancement of the level of 2',5'-oligo-adenylate synthetase (2',5'-A); and (6) enhancement of the double-stranded RNA-dependent protein kinase. The sonicates were active in inhibiting viral infection in both human and other mammalian cells such as hamster, monkey, mouse, and rabbit cells.

The tests show that IFN- $\alpha$ 54 exhibits anti-10 viral activity against DNA and RNA viruses, cell growth regulating activity, and an ability to regulate the production of intracellular enzymes and other cell-produced substances. Accordingly, it is expected IFN-c54 may be used to treat viral infections with a 15 potential for interferon therapy such as chronic hepatitis B infection, ocular, local, or systemic herpes virus infections, influenza and other respiratory tract virus infections, rabies and other viral zoonoses, arbovirus infections, and slow virus diseases 20 such as Kuru and sclerosing panencephalitis. It may also be useful for treating viral infections in immunocompromised patients such as herpes zoster and varicella, cytomegalovirus, Epstein-Barr virus infection, herpes simplex infections, rubella, and progres-25 sive multifocal leukoencephalopathy. Its cell growth regulating activity makes it potentially useful for treating tumors and cancers such as osteogenic sarcoma, multiple myeloma, Hodgkin's disease, nodular, poorly differentiated lymphoma, acute lymphocytic 30 leukemia, breast carcinoma, melanoma, and nasopharyngeal carcinoma. The fact that IFN- $\alpha$ 54 increases protein kinase and 2',5'-oligoadenylate synthetase



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indicates it may also increase synthesis of other enzymes or cell-produced substances commonly affected by IFNs such as histamine, hyaluronic acid, prostaglandin E, ItRNA methylase, and aryl hydrocarbon hydro-5 lase. Similarly, it may be useful to inhibit enzymes commonly inhibited by IFNs such as tyrosine amino transferase, glycerol-3-phosphate dehydrogenase glutamine synthetase, ornithine decarboxylase, S-adenosyl-1-methionine decarboxylase, and UDP-N-10 acetylglucosamine-dolichol monophosphate transferase. The ability of the IFN- $\alpha$ 54 to stimulate NK cell activity is indicative that it may also possess other activities such as the abilities to induce macrophage activity and antibody production and to effect cell 15 surface alterations such as changes in plasma membrane density or cell surface charge, altered capacity to bind substances such as cholera toxin, concanavalin A and thyroid-stimulating hormone, and change in the exposure of surface gangliosides.

Pharmaceutical compositions that contain IFN-α54 as an active ingredient will normally be formulated with an appropriate solid or liquid carrier depending upon the particular mode of administration being used. For instance, parenteral formulations are usually injectable fluids that use pharmaceutically and physiologically acceptable fluids such as physiological saline, balanced salt solutions, or the like as a vehicle. Oral formulations, on the other hand, may be solid, eg tablet or capsule, or liquid solutions or suspensions. IFN-α54 will usually be formulated as a unit dosage form that contains in the range of 10<sup>4</sup> to 10<sup>7</sup> international units, more usually 10<sup>6</sup> to 10<sup>7</sup> international units, per dose.



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IFN- $\alpha$ 54 may be administered to humans in various manners such as orally, intravenously, intramuscularly, intraperitoneally, intranasally, intradermally, and subcutaneously. The particular mode of 5 administration and dosage regimen will be selected by the attending physician taking into account the particulars of the patient, the disease and the disease state involved. For instance, viral infections are usually treated by daily or twice daily doses over a 10 few days to a few weeks; whereas tumor or cancer treatment involves daily or multidaily doses over months or years. IFN- $\alpha$ 54 therapy may be combined with other treatments and may be combined with or used in association with other chemotherapeutic or chemo-15 preventive agents for providing therapy against viral infections, neoplasms, or other conditions against which it is effective. For instance, in the case of herpes virus keratitis treatment, therapy with IFN has been supplemented by thermocautery, debridement and 20 trifluorothymidine therapy.

Modifications of the above described modes for carrying out the invention, such as, without limitation, use of alternative vectors, alternative expression control systems in the vector, and alternative host microorganisms and other therapeutic or related uses of IFN-α54, that are obvious to those of ordinary skill in the biotechnology, pharmaceutical, medical and/or related fields are intended to be within the scope of the following claims.



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#### Claims

1. A polypeptide having interferon activity and comprising the amino acid sequence:

CysAspLeuProGln ThrHisSerLeuGly HisArgArgThrMet MetLeuLeuAlaGln MetArgArgIleSer LeuPheSerCysLeu LysAspArgHisAsp PheArgPheProGln GluGluPheAspGly AsnGlnPheGlnLys AlaGluAlaIleSer ValLeuHisGluVal IleGlnGlnThrPhe AsnLeuPheSerThr LysAspSerSerVal AlaTrpAspGluArg LeuLeuAspLysLeu TyrThrGluLeuTyr GlnGlnLeuAsnAsp LeuGluAlaCysVal MetGlnGluValTrp ValGlyGlyThrPro LeuMetAsnGluAsp SerIleLeuAlaVal ArgLysTyrPheGln ArgIleThrLeuTyr LeuThrGluLysLys TyrSerProCysAla ArgLeuArgArgLys Glu.

- 2. The polypeptide of claim 1 wherein the polypeptide consists essentially of said amino acid sequence.
- 3. The polypeptide of claim 1 or 2 wherein the initial cysteine residue of the amino acid sequence is preceded by an N-formyl-methionine group.
  - 4. The polypeptide of claim 1 or 2 wherein the amino acid residues of said sequence are unsubstituted.
    - 5. IFN- $\alpha$ 54.
- 6. A composition having interferon activity and comprising a mixture of:
  - (a) a polypeptide having the amino acid sequence

CysAspLeuProGln ThrHisSerLeuGly HisArgArgThrMet MetLeuLeuAlaGln MetArgArgIleSer LeuPheSerCysLeu LysAspArgHisAsp PheArgPheProGln GluGluPheAspGly AsnGlnPheGlnLys AlaGluAlaIleSer ValLeuHisGluVal IleGlnGlnThrPhe AsnLeuPheSerThr LysAspSerSerVal AlaTrpAspGluArg LeuLeuAspLysLeu TyrThrGluLeuTyr GlnGlnLeuAsnAsp LeuGluAlaCysVal



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MetGlnGluValTrp ValGlyGlyThrPro LeuMetAsnGluAsp SerIleLeuAlaVal ArgLysTyrPheGln ArgIleThrLeuTyr LeuThrGluLysLys TyrSerProCysAla TrpGluValValArg AlaGluIleMetArg SerPheSerSerSer ArgAsnLeuGlnGlu ArgLeuArgArgLys Glu

and;

- (b) a polypeptide having said amino acid sequence wherein the initial cysteine residue of the sequence is preceded by an N-formyl-methionine or methionine group.
  - 7. The composition of claim 6 wherein the amino acid residues of said sequence are unsubstituted.
- 10 8. A composition having interferon activity comprising a polypeptide having the amino acid sequence

CysAspLeuProGln ThrHisSerLeuGly HisArgArgThrMet MetLeuLeuAlaGln MetArgArgIleSer LeuPheSerCysLeu LysAspArgHisAsp PheArgPheProGln GluGluPheAspGly AsnGlnPheGlnLys AlaGluAlaIleSer ValLeuHisGluVal IleGlnGlnThrPhe AsnLeuPheSerThr LysAspSerSerVal AlaTrpAspGluArg LeuLeuAspLysLeu TyrThrGluLeuTyr GlnGlnLeuAsnAsp LeuGluAlaCysVal MetGlnGluValTrp ValGlyGlyThrPro LeuMetAsnGluAsp SerIleLeuAlaVal ArgLysTyrPheGln ArgIleThrLeuTyr LeuThrGluLysLys TyrSerProCysAla TrpGluValValArg AlaGluIleMetArg SerPheSerSerSer ArgAsnLeuGlnGlu ArgLeuArgArgLys Glu

- or a mixture of said polypeptide and a polypeptide

  15 having said sequence wherein the initial cysteine
  residue is preceded by an N-formyl-methionine or
  methionine group wherein the interferon activity of
  the composition is attributable to said polypeptide or
  to said mixture.
- 9. A DNA unit consisting of a nucleotide sequence that encodes the polypeptide of claim 1 or 5.



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10. The DNA unit of claim 9 wherein the nucleotide sequence is:

TGTGATCTGCCTCAGACCCACAGCCTGGGTCACAGGAG

- 11. A cloning vehicle that includes the DNA unit of claim 9 or 10.
  - 12. The cloning vehicle of claim 11 wherein the cloning vehicle is a plasmid.
  - 13. The cloning vehicle of claim 11 wherein the cloning vehicle is the plasmid pCS12.
- 10 14. A host that is transformed with the cloning vehicle of claim 11 and produces IFN- $\alpha$ 54.
  - 15. The host of claim 14 wherein the host is a prokanyote.

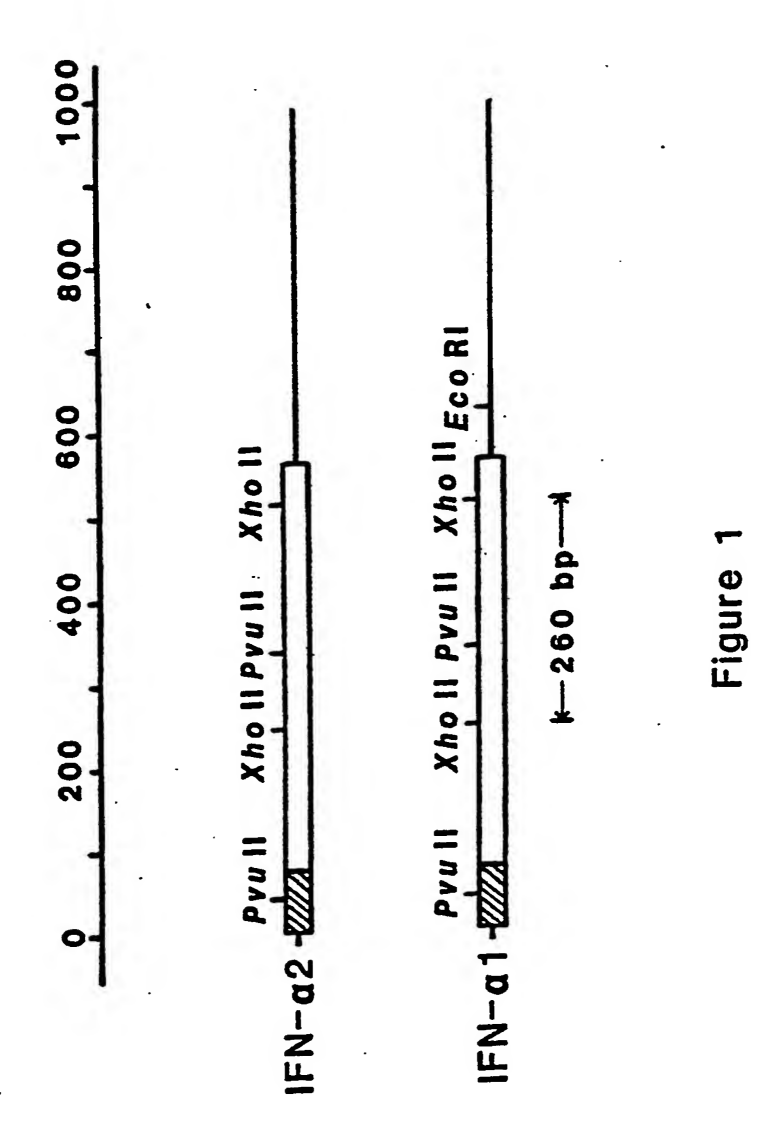


- 16. The host of claim 14 wherein the host organism is E.coli.
- 17. A host that is transformed with the cloning vehicle of claim 13 and produces IFN- $\alpha$ 54, wherein the host is <u>E.coli</u>.
  - 18. A process for producing IFN- $\alpha$ 54 comprising cultivating the host of claim 14 and collecting IFN- $\alpha$ 54, from the resulting culture.
- 19. A process of producing IFN-α54 compri-10 sing cultivating the host organism of claim 16 and collecting IFN-α54 from the resulting culture.
  - 20. A process for producing IFN- $\alpha$ 54 comprising cultivating the host organism of claim 17 and collecting IFN- $\alpha$ 54 from the resulting culture.
- 21. A pharmaceutical composition comprising an effective amount of the polypeptide of claim 1, 2 or 5 admixed with a pharmaceutically acceptable vehicle or carrier.
- 22. A pharmaceutical composition comprising an effective amount of the composition of claim 6 or 8 admixed with a pharmaceutically acceptable vehicle or carrier.
- 23. A method of providing interferon therapy to a human comprising administering a therapeutically effective amount of the polypeptide of claim 1, 2 or 5 to said human.

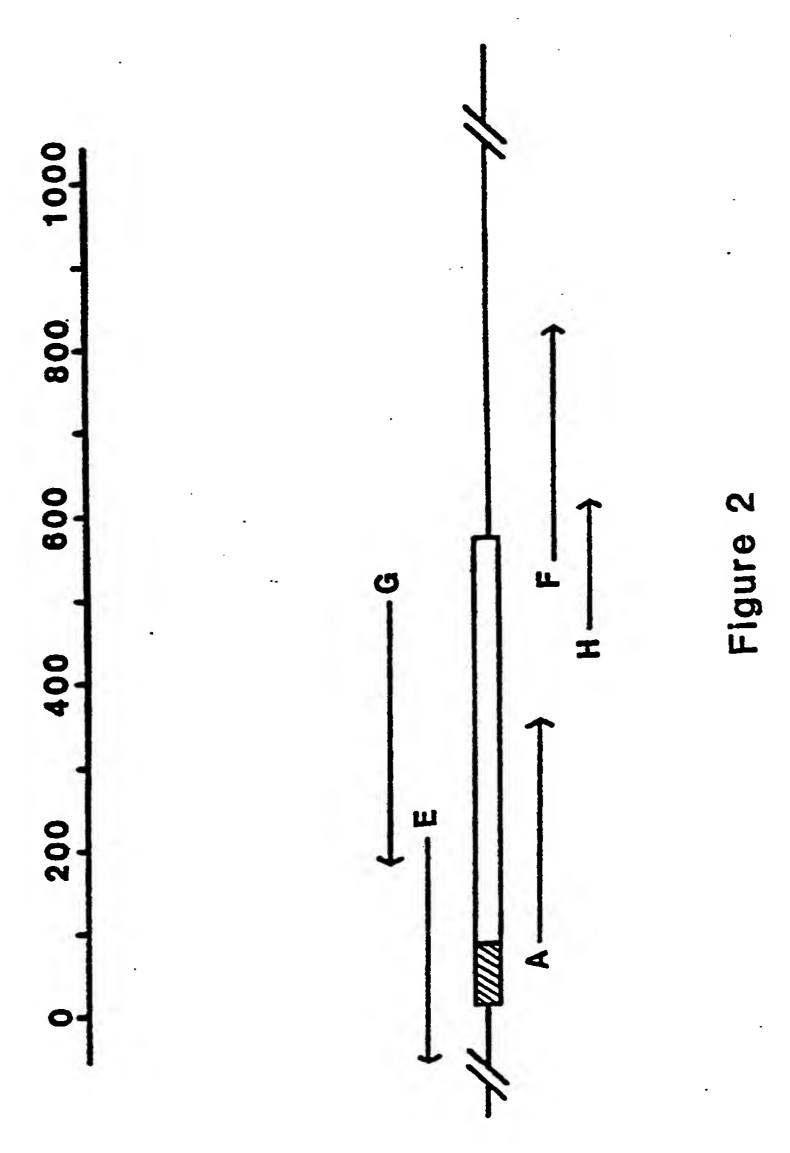


- 24. A method of providing interferon therapy to a human comprising administering a therapeutically effective amount of the composition of claim 6 or 8 to said human.
- 25. The method of claim 23 wherein the therapy is for treating a viral infection, providing cell growth regulation, or regulating the production of a cell-produced substance.
- 26. The method of claim 24 wherein the 10 therapy is for treating a viral infection, providing cell growth regulation, or regulating the production of a cell-produced substance.
- 27. A method of providing antiviral therapy to a mammal comprising administering a viral infection inhibiting amount of the polypeptide of claim 1, 2 or 5 to the mammal.











		30	40	. 50	- 60
10	20.		ACAAGTCCAA		AACATCTACA
	•		TGTTCAGGTT	GTCGTAGACG	TTGTAGATGT
TCTCGACTTC	<b></b>		100	110	120
70	80	90	•		GTCAAGCTGC
ATGGCTTTGC		ACTGATGGCC	GACCACCACG		CAGTTCGACG
TACCGAAACG	<b>4.234</b>	TGACTACCGG	160	170	180
130	140	150	AGCCTGGGTC		CATGATGCTC
TCTCTGGACT	GTGATCTGCC		TCGGACCCAG	TGTCCTCCTG	GTACTACGAG
AGAGACCTGA	CACTAGACGG	AGTCTGGGTG	220	230	240
190	200	210		AGGACAGACA	TGACTTCAGA
CTGGCACAAA	TGAGGAGAAT		AGGACAGACT	TCCTGTCTGT	ACTGAAGTCT
GACCGTGTTT	ACTCCTCTTA		280	290	300
250-	260	270			CTCTGTCCTC
TTTCCCCAGG	AGGAGTTTGA		AAGGTCTTCC	GACTTCGGTA	
AAAGGGGTCC	TCCTCAAACT	ACCGTTGGTC	340	350	360
310	320	330		AGGACTCATC	TGTTGCTTGG
CATGAGGTGA		CTTCAATCTC	=	TCCTGAGTAG	ACAACGAACC
GTACTCCACT	AAGTCGTCTG	GAAGTTAGAG	AAGTCGTGTT 400	410	420
. 370	. 380	390		AGCAGCTGAA	TGACCTGGAA
GATGAGAGGC	TTCTAGACAA	•	GAACTTTACC	TCGTCGACTT	ACTGGACCTT
CTACTCTCCG	AAGATCTGTT	TGAGATATGA	CTTGAAATGG	470	480
430	440	450	460	TGATGAATGA	
GCCTGTGTGA	TGCAGGAGGT	GTGGGTGGGA	GGGACTCCCC	ACTACTACT	CCTGAGGTAG
CGGACACACT	ACGTCCTCCA		CCCTGAGGGG	530	540
490	500	510	520		
CTGGCTGTGA				TGACAGAGAA	TTTCATGTCG
GACCGACACT	<del>-</del> - ··	GGTTTCTTAG		ACTGTCTCTT 590	600
550	560	570	580		
CCTTGTGCCT	GGGAGGTTGT	CAGAGCAGAA	ATCATGAGAT	CCTTCTCTTC	TAGTTCTTTG
GGAACACGGA	<del>-</del>	GTCTCGTCTT		GGAAGAGAAG 650	660
610	620	630	,640		<del>-</del>
TTGCAAGAAA	GGTTAAGGAG	GAAGGAATAA	GACCTGATCC	AACACAGAAA	CCTCAGGGTA
AACGTTCTTT	CCAATTCCTC		CTGGACTAGG	TTGTGTCTTT 710	720
670	680	690	700		• — -
TGACGACTAC	ACCAGCTTGC	ACTTTCATGA	TCTGCCATTT	TAAAGACTCT	ACABAGACGA
ACTGCTGATG		TGAAAGTACT	AGACGGTAAA	ATTTCTGAGA 770	780
730	740	750			TAACGAACAT
ATAACCATAC		•	TCAAGTATTT	AGTTCACACA	
TATTGGTATG				830	*** * * * * * * * * * * * * * * * * * *
790	800	.810	820	•	ATCTATTT
CGTGTTCAGT				AACGTGATGG TTGCACTACC	TAGATAAA
GCACAAGTCA	ACGTGTCCTT	GATCAGGGAA	TGTCTACTGA	TIGNOTAGE	<b>→ = = → = →</b> → → → → → → → → → → → → → →

Figure 3



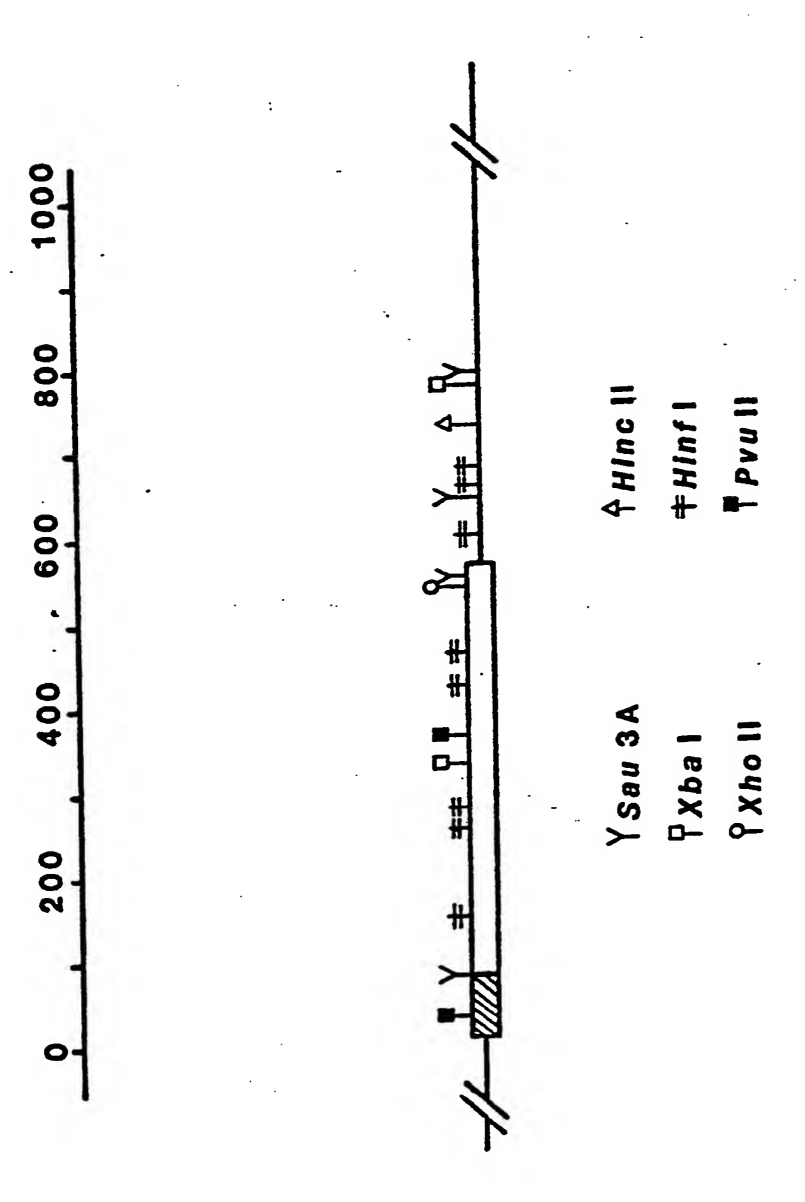




Figure 4

Met Ala Leu Pro Phe Ala Leu Leu Het Ala Leu Val Val Leu Ser Cys Lys Ser Ser Cys ATG GCT TTG CCT TTT GCT TTA CTG ATG GCC CTG GTG GTG CTC AGC TGC AAG TCA AGC TGC Ser Leu Asp Cys Asp Leu Pro Gln Thr His Ser Leu Gly His Arg Arg Thr Met Het Leu TCT CTG GAC TGT GAT CTG CCT CAG ACC CAC AGC CTG GGT CAC AGG AGG ACC ATG ATG CTC Leu Ala Gln Met Arg Arg Ile Ser Leu Phe Ser Cys Leu Lys Asp Arg His Asp Phe Arg CTG GCA CAA ATG AGG AGA ATC TCT CTT TTC TCC TGT CTG AAG GAC AGA CAT GAC TTC AGA Phe Pro Gln Glu Glu Phe Asp Gly Asn Gln Phe Gln Lys Ala Glu Ala Ile Ser Val Leu TTT CCC CAG GAG GAG TTT GAT GGC AAC CAG TTC CAG AAG GCT GAA GCC ATC TCT GTC CTC His Glu Val Ile Gln Gln Thr Phe Asn Leu Phe Ser Thr Lys Asp Ser Ser Val Ala Trp CAT GAG GTG ATT CAG CAG ACC TTC AAT CTC TTC AGC ACA AAG GAC TCA TCT GTT GCT TGG Asp Glu Arg Leu Leu Asp Lys Leu Tyr Thr Glu Leu Tyr Gln Gln Leu Asn Asp Leu Glu GAT GAG AGG CTT CTA GAC AAA CTC TAT ACT GAA CTT TAC CAG CAG CTG AAT GAC CTG GAA Ala Cys Val Met Gln Glu Val Trp Val Gly Gly Thr Pro Leu Met Asn Glu Asp Ser Ile GCC TGT GTG ATG CAG GAG GTG GTG GGA GGG ACT CCC CTG ATG AAT GAG GAC TCC ATC Leu Ala Val Arg Lys Tyr Phe Gln Arg Ile Thr Leu Tyr Leu Thr Glu Lys Lys Tyr Ser CTG GCT GTG AGA AAA TAC TTC CAA AGA ATC ACT CTC TAC CTG ACA GAG AAA AAG TAC AGC Pro Cys Ala Trp Glu Val Val Arg Ala Glu Ile Net Arg Ser Phe Ser Ser Ser Arg Asn CCT TGT GCC TGG GAG GTT GTC AGA GCA GAA ATC ATG AGA TCC TTC TCT TCA TCA AGA AAC Leu Gln Glu Arg Leu Arg Arg Lys Glu TTG CAA GAA AGG TTA AGG AGG AAG GAA 181

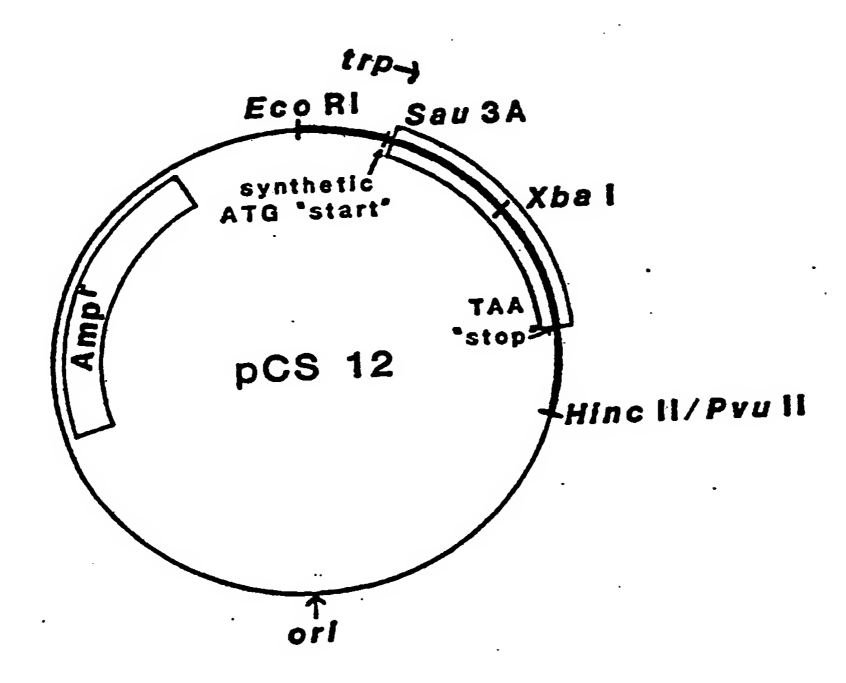
Figure 5



GAA TTC, CGA CAT CAT AAC GGT TCT GGC AAA TAT TCT GAA ATG AGC TGT TGA CAA TTA ATC Met Cys Eco RI ATC GAA CTA GTT AAC TAG TAC GCA AGT TCA CGT AAA AAG GGT ATC GAT AAG CTT ATG TGT Asp Leu Pro Gln Thr His Ser Leu Gly His Arg Arg Thr Met Met Leu Leu Ala Gln Met GAT CTG CCT CAG ACC CAC AGC CTG GGT CAC AGG AGG ACC ATG ATG CTC CTG GCA CAA ATG Arg Arg Ile Ser Leu Phe Ser Cys Leu Lys Asp Arg His Asp Phe Arg Phe Pro Gln Glu AGG AGA ATC TCT CTT TTC TCC TGT CTG AAG GAC AGA CAT GAC TTC AGA TTT CCC CAG GAG Glu Phe Asp Gly Asn Gln Phe Gln Lys Ala Glu Ala Ile Ser Val Leu His Glu Val Ile GAG TTT GAT GGC AAC CAG TTC CAG AAG GCT GAA GCC ATC TCT GTC CTC CAT GAG GTG ATT Gln Gln Thr Phe Asn Leu Phe Ser Thr Lys Asp Ser Ser Val Ala Trp Asp Glu Arg Leu CAG CAG ACC TTC AAT CTC TTC AGC ACA AAG GAC TCA TCT GTT GCT TGG GAT GAG AGG CTT Leu Asp Lys Leu Tyr Thr Glu Leu Tyr Gln Gln Leu Asn Asp Leu Glu Ala Cys Val Met CTA GAC AAA CTC TAT ACT GAA CTT TAC CAG CAG CTG AAT GAC CTG GAA GCC TGT GTG ATG Gln Glu Val Trp Val Gly Gly Thr Pro Leu Met Asn Glu Asp Ser Ile Leu Ala Val Arg Xba I CAG GAG GTG TGG GTG GGG ACT CCC CTG ATG AAT GAG GAC TCC ATC CTG GCT GTG AGA Lys Tyr Phe Gln Arg Ile Thr Leu Tyr Leu Thr Glu Lys Lys Tyr Ser Pro Cys Ala Trp AÃA TÁC TTC CAA AGA ATC ACT CTC TÁC CTG ACA GAG AÃA AÃG TÁC AGC CCT TGT GCC TGG Glu Val Val Arg Ala Glu Ile Met Arg Ser Phe Ser Ser Ser Arg Asn Leu Gln Glu Arg GAG GTT GTC AGA GCA GAA ATC ATG AGA TCC TTC TCT TCA TCA AGA AAC TTG CAA GAA AGG 601 TTA AGG AGG AAG GAA TAA GAC CTG ATC CAA CAC AGA AAC GAC TCC CAT TGA CGA CTA CAC CAG CTT GCA CTT TCA TGA TCT GCC ATT TTA AAG ACT CTT GTT TCT GCT ATA ACC ATA CCA TGA GTT GAA TCA AAC GCG TCA AGT ATT TTC AAG TGT GTT

Figure 6





IFN-a54 Expression Plasmid

Figure 7



## INTERNATIONAL SEARCH REPORT\_

International Application No PCT/US 83/00033

I. CLAS	SIFICATION OF SUBJECT MATTER (If severa	al classification symbols apply, indicate all) s	
According	g to International Patent Classification (IPC) or to b	oth National Classification and IPC	N 61 ₹ 45/02-
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IIL DOCI	UMENTS CONSIDERED TO BE RELEVANT	•	Relevant to Claim No. 18
Category *	Citation of Document, 16 with Indication, wh	ere appropriate, of the relevant passages 17	Keisasut m Cramii Mor
Y	Nature, volume 290, 5 Mature, volume 290, 5 Mature, D.Goeddel et al.: eight distinct cloral interferon C DNA's see the entire documents.	The structure of ned human leukocyte , pages 20-26,	1,4,8-12
<b>Y</b>	Nature, volume 287, 2 of D.Goeddel et al.: 'interferon produced biologically active see the entire documents of the contraction of the application of the application.	'Human leukocyte d by E.Coli is e", pages 411-416, ment	1,4,8-12
Y	Proc.Natl.Acad.Sci, vol September 1981 (US) major human leukoc gene", pages 5435-5 see the entire docu	"DNA sequence of a cyte interferon 439,	1,4,8-10
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E	UROPEAN PATENT OFFICE	G.L.M	kruydenberg

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Y	Science, volume 209, 19 September 1980, M.Streuli et al.: "At least three human type alpha interferons: Structure of alpha 2", pages 1343-1347, see the entire document (cited in the application)	1,4,8-10
¥	EP, A, 0042246 (CANCER INSTITUTE OF JAPANESE FOUNDATION FOR CANCER RESEARCH) 23 December 1981, see claims 1-8	1,2,4,8-12
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